

RAW SEQUENCE LISTING ERROR REPORT

The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable form:

Application Serial Number:	10/019,151
Source:	05/10
Date Processed by STIC:	3/19/2003
· ·	

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.
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FOR CRF SUBMISSION QUESTIONS, PLEASE CONTACT MARK SPENCER, 703-308-4212.

FOR SEQUENCE RULES INTERPRETATION, PLEASE CONTACT ROBERT WAX, 703-308-4216. PATENTIN 2.1 e-mail help: patin21help@uspto.gov or phone 703-306-4119 (R. Wax) PATENTIN 3.0 e-mail help: patin3help@uspto.gov or phone 703-306-4119 (R. Wax)

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE CHECKER VERSION 3.1 PROGRAM, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW FOR ADDRESS:

http://www.uspto.gov/web/offices/pac/checker

Applicants submitting genetic sequence information electronically on diskette or CD-Rom should be aware that there is a possibility that the disk/CD-Rom may have been affected by treatment given to all incoming mail.

Please consider using alternate methods of submission for the disk/CD-Rom or replacement disk/CD-Rom.

Any reply including a sequence listing in electronic form should NOT be sent to the 20231 zip code address for the United States Patent and Trademark Office, and instead should be sent via the following to the indicated addresses:

- 1. EFS-Bio (httm, EFS Submission User Manual ePAVE)
- 2. U.S. Postal Service: U.S. Patent and Trademark Office, Box Sequence, P.O. Box 2327, Arlington, VA 22202
- Hand Carry directly to:
 U.S. Patent and Trademark Office, Technology Center 1600, Reception Area, 7th Floor, Examiner Name, Sequence Information, Crystal Mall One, 1911 South Clark Street, Arlington, VA 22202
 - U.S. Patent and Trademark Office, Box Sequence, Customer Window, Lobby, Room 1B03, Crystal Plaza Two, 2011 South Clark Place, Arlington, VA 22202
- 4. Federal Express, United Parcel Service, or other delivery service to: U.S. Patent and Trademark Office, Box Sequence, Room 1B03-Mailroom, Crystal Plaza Two, 2011 South Clark Place, Arlington, VA 22202

Revised 01/29/2002

ERROR DETECTED	suggested correction serial number: 10/0/9/5	
attn: New Rules Case	s: Please disregard english "Alpha" headers, which yere inserted by Pto S	OFTWARE
lWrapped Nucleics Wrapped Aminos	The number/text at the end of each line "wrapped" down to the next line. This may occur if your file was retrieved in a word processor after creating it. Please adjust your right margin to .3; this will prevent "wrapping."	
2Invalid Line Length	The rules require that a line not exceed 72 characters in length. This includes white spaces.	-
Misaligned Amino Numbering	The numbering under each 5th amino acid is misaligned. Do not use tab codes between numbers; the use space characters, instead.	
4Non-ASCII	The submitted file was not saved in ASCII(DOS) text, as required by the Sequence Rules. Please ensure your subsequent submission is saved in ASCII text.	
5Variable Length	Sequence(s)contain n's or Xaa's representing more than one residue. Per Sequence Rules, each n or Xaa can only represent a single residue. Please present the maximum number of each residue having variable length and indicate in the <220>-<223> section that some may be missing.	
6Patentin 2.0 "bug"	A "bug" in PatentIn version 2.0 has caused the <220>-<223> section to be missing from amino acid sequences(s) Normally, PatentIn would automatically generate this section from the previously coded nucleic acid sequence. Please manually copy the relevant <220>-<223> section to the subsequent amino acid sequence. This applies to the mandatory <220>-<223> sections for Artificial or Unions win sequences.	
7Skipped Sequences (OLD RULES)	Sequence(s) missing. If intentional, please insert the following lines for each skipped sequence: (2) INFORMATION FOR SEQ ID NO:X: (insert SEQ ID NO where "X" is shown) (i) SEQUENCE CHARACTERISTICS: (Do not insert any subheadings under this heading) (xi) SEQUENCE DESCRIPTION:SEQ ID NO:X: (insert SEQ ID NO where "X" is shown) This sequence is intentionally skipped	
	Please also adjust the "(ii) NUMBER OF SEQUENCES:" response to Include the skipped sequences.	
8Skipped Sequences (NEW RULES)	Sequence(s) missing. If Intentional, please insert the following lines for each skipped sequence. <210> sequence id number <400> sequence id number 000	
9Use of n's or Xaa's (NEW RULES)	Use of n's and/or Xaa's have been detected in the Sequence Listing. Per 1.823 of Sequence Rules, use of <220>-<223> is MANDATORY if n's or Xaa's are present. In <220> to <223> section, please explain location of n or Xaa; and which residue n or Xaa represents.	·
0Invalid <213> Response	Per 1.823 of Sequence Rules, the only valld <213> responses are: Unknown, Artificial Sequence, or scientific name (Genuz/species). <220> <223> section is required when <213> response is Unknown or sis Artificial Sequence	nus.
1Use of <220>	Sequence(s) missing the <220> "Feature" and associated numeric identifiers and responses. Use of <220> to <223> is MANDATORY if <213> "Organism" response is "Artificial Sequence" or "Unknown." Please explain source of genetic material in <220> to <223> section. (See "Federal Register," 06/01/1998, Vol. 63, No. 104, pp. 29631-32) (Sec. 1.823 of Sequence Rules)	
2Patentin 2.0 "bug"	Please do not use "Copy to Disk" function of Patentin version 2.0. This causes a corrupted file, resulting in missing mandatory numeric identifiers and responses (as indicated on raw sequence listing). Instead, please use "File Manager" or any other manual means to copy file to floppy disk.	•
3Misuse of n	n can only be used to represent a single nucleotide in a nucleic acid sequence. N is not used to represent any value not specifically a nucleotide.	

AMC/MH - Biotechnology Systems Branch - 08/21/2001



DATE: 03/19/2003 RAW SEQUENCE LISTING PATENT APPLICATION: US/10/019,151 TIME: 14:33:58

Input Set: A:\(01997.025700) Sequence Listing.txt Output Set: N:\CRF4\03192003\J019151.raw

4 <110> APPLICANT: Sagami Chemical Research Center, Protegene Inc.

6 <120> TITLE OF INVENTION: Human proteins having hydrophobic domains and DNAs encoding

these proteins

9 <130> FILE REFERENCE: 661924

11 <140> CURRENT APPLICATION NUMBER: US/10/019,151

11 <141> CURRENT FILING DATE: 2003-01-31

11 <150> PRIOR APPLICATION NUMBER: JP 11-178065

12 <151> PRIOR FILING DATE: 1999-06-24

14 <160> NUMBER OF SEQ ID NOS: 24

pp 1-11 evere shoughout Does Not Comply ERRORED SEQUENCES Corrected Diskette Needed 16 <210> SEQ ID NO: 1

17 <211> LENGTH: 238

18 <212> TYPE: PRT

19 <213> ORGANISM: Homo sapiens

21 <400> SEQUENCE: 1

Met Ile Leu Leu Val Ile Leu Ala Phe Tyr Leu Trp Gln Val Asp Met 23 Leu Ser Glu Ile Asn Ile Ala Pro Arg Ile Leu Thr Asn Phe Thr Gly 24 25 25 26 Val Met Pro Pro Gln Phe Lys Lys Asp Leu Asp Ser Tyr Leu Lys Thr E--> 27 28 Arg Ser Pro Val Thr Phe Leu Ser Asp Leu Arg Ser Asn Leu Gln Val E--> 29 55 60

Ser Asn Glu Pro Gly Asn Arg Tyr Asn Leu Gln Leu Ile Asn Ala Leu 30 E --> 31Val Leu Tyr Val Gly Thr Gln Ala Ile Ala His Ile His Asn Lys Gly 32

E--> 33 90 Ser Thr Pro Ser Met Ser Thr Ile Thr His Ser Ala His Met Asp Ile 34

E--> 35 105 Phe Gln Asn Leu Ala Val Asp Leu Asp Thr Glu Gly Arg Tyr Leu Phe 36

E--> 37 120 38 Leu Asn Ala Ile Ala Asn Gln Leu Arg Tyr Pro Asn Ser His Thr His E--> 39

135 140 Tyr Phe Ser Cys Thr Met Leu Tyr Leu Phe Ala Glu Ala Asn Thr Glu 40 E--> 41 150 155

42 Ala Ile Gln Glu Gln Ile Thr Arg Val Leu Leu Glu Arg Leu Ile Val E --> 43165 170

Asn Arg Pro His Pro Trp Gly Leu Leu Ile Thr Phe Ile Glu Leu Ile 44 E --> 45185

Lys Asn Pro Ala Phe Lys Phe Trp Asn His Glu Phe Val His Cys Ala

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anio acid
humbus
1 sel etem 3
on Eva Summary
Meet)

Input Set : A:\(01997.025700) Sequence Listing.txt
Output Set: N:\CRF4\03192003\J019151.raw

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E--> 47
                  195
                                      200
          Pro Glu Ile Glu Lys Leu Phe Gln Ser Val Ala Gln Cys Cys Met Gly
     48
                                  215
 --> 49
          Gln Lys Gln Ala Gln Gln Val Met Glu Gly Thr Gly Ala Ser
     50
E--> 51
                              230
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     54 <211> LENGTH: 339
     55 <212> TYPE: PRT
     56 <213> ORGANISM: Homo sapiens
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          Gly Leu His Leu Phe Leu Leu Thr Ala Gly Pro Ala Leu Gly Trp Asn
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     62
     63
          Asp Pro Asp Arg Met Leu Leu Arg Asp Val Lys Ala Leu Thr Leu His
     64
     65
          Tyr Asp Arg Tyr Thr Thr Ser Arg Arg Leu Asp Pro Ile Pro Gln Leu
     66
          Lys Cys Val Gly Gly Thr Ala Gly Cys Asp Ser Tyr Thr Pro Lys Val
     67
     68
                               70
     69
          Ile Gln Cys Gln Asn Lys Gly Trp Asp Gly Tyr Asp Val Gln Trp Glu
     70
                           8.5
     71
          Cys Lys Thr Asp Leu Asp Ile Ala Tyr Lys Phe Gly Lys Thr Val Val
     72
                                          105
     73
          Ser Cys Glu Gly Tyr Glu Ser Ser Glu Asp Gln Tyr Val Leu Arg Gly
     74
                                      120
     75
          Ser Cys Gly Leu Glu Tyr Asn Leu Asp Tyr Thr Glu Leu Gly Leu Gln
     77
          Lys Leu Lys Glu Ser Gly Lys Gln His Gly Phe Ala Ser Phe Ser Asp
     78
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                                                  155
     79
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     80
                          165
                                              170
     81
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E--> 82
                      180 180
                                          185
                                                       · 190
     83
          Phe Leu Ser Asp Gly Gln Tyr Ser Pro Pro Pro Tyr Ser Glu Tyr Pro
E--> 84
                                      200
                                                           205
    85
          Pro Phe Ser His Arg Tyr Gln Arg Phe Thr Asn Ser Ala Gly Pro Pro
E--> 86
                                  215
     87
          Pro Pro Gly Phe Lys Ser Glu Phe Thr Gly Pro Gln Asn Thr Gly His
E--> 88
                              230
                                                  235
          Gly Ala Thr Ser Gly Phe Gly Ser Ala Phe Thr Gly Gln Gln Gly Tyr
     89
E --> 90
          Glu Asn Ser Gly Pro Gly Phe Trp Thr Gly Leu Gly Thr Gly Gly Ile
     91
E--> 92
                                           265
     93
          Leu Gly Tyr Leu Phe Gly Ser Asn Arg Ala Ala Thr Pro Phe Ser Asp
E--> 94
                                      280
     95
          Ser Trp Tyr Tyr Pro Ser Tyr Pro Pro Ser Tyr Pro Gly Thr Trp Asn
E--> 96
                                  295
     97
          Arg Ala Tyr Ser Pro Leu His Gly Gly Ser Gly Ser Tyr Ser Val Cys
```

misoligned, humbering

Input Set : A:\(01997.025700) Sequence Listing.txt

Output Set: N:\CRF4\03192003\J019151.raw

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325
330

Arg Arg Arg
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E--> 100
     101
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     111
           Leu Ser Ala Ile Leu Ser Met Leu Ser Leu Ser Phe Ser Thr Thr Ser
     112
     113
           Leu Leu Ser Asn Tyr Trp Phe Val Gly Thr Gln Lys Val Pro Lys Pro
     114
                                        40
     115
           Leu Cys Glu Lys Gly Leu Ala Ala Lys Cys Phe Asp Met Pro Val Ser
     116
                                    55
     117
           Leu Asp Gly Asp Thr Asn Thr Ser Thr Gln Glu Val Val Gln Tyr Asn
     118
                                70
           Trp Glu Thr Gly Asp Asp Arg Phe Ser Phe Arg Ser Phe Arg Ser Gly
     119
     120
     121
           Met Trp Leu Ser Cys Glu Glu Thr Val Glu Glu Pro Gly Glu Arg Cys
     122
                                           105
     123
           Arg Ser Phe Ile Glu Leu Thr Pro Pro Ala Lys Arg Glu Ile Leu Trp
     124
                                       120
     125
           Leu Ser Leu Gly Thr Gln Ile Thr Tyr Ile Gly Leu Gln Phe Ile Ser
     126
                                   135
     127
           Phe Leu Leu Leu Thr Asp Leu Leu Thr Gly Asn Pro Ala Cys
     128
                               150
                                                   155
     129
           Gly Leu Lys Leu Ser Ala Phe Ala Ala Val Ser Ser Val Leu Ser Gly
     130
                           165
                                               170
     131
           Leu Leu Gly Met Val Ala His Met Met Tyr Ser Gln Val Phe Gln Ala
     132
                                           185
     133
           Thr Val Asn Leu Gly Pro Glu Asp Trp Arg Pro His Val Trp Asn Tyr
     134
                                       200
     135
           Gly Trp Ala Phe Tyr Met Ala Trp Leu Ser Phe Thr Cys Cys Met Ala
     136
                                   215
                                                       220
     137
           Ser Ala Val Thr Thr Phe Asn Thr Tyr Thr Arg Met Val Leu Glu Phe
     138
                               230
                                                   235
     139
           Lys Cys Lys His Ser Lys Ser Phe Lys Glu Asn Pro Asn Cys Leu Pro
     140
                           245
                                               250
     141
           His His Gln Cys Phe Pro Arg Arg Leu Ser Ser Ala Ala Pro Thr
     142
                                           265
     143
           Val Gly Pro Leu Thr Ser Tyr His Gln Tyr His Asn Gln Pro Ile His
     144
                                       280
           Ser Val Ser Glu Gly Val Asp Phe Tyr Ser Glu Leu Arg Asn Lys Gly
     145
     146
                                   295
E--> 147
           Phe Gin Arg Gly Ala Ser Gln Glu Leu Lys Glu Ala Val Arg Ser Ser
     148
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Input Set : A:\(01997.025700) Sequence Listing.txt
Output Set: N:\CRF4\03192003\J019151.raw

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     207
                                                 1.0
     208
           Met Asn Ser Arg Gly Ile Trp Leu Ser Tyr Val Leu Ala Ile Gly Leu
     209
                        20
                                             25
           Leu His Ile Val Leu Leu Ser Ile Pro Phe Val Ser Val Pro Val Val
     210
     211
                                         40
           Trp Thr Leu Thr Asn Leu Ile His Asn Met Gly Met Tyr Ile Phe Leu
     212
     213
                                     55
           His Thr Val Lys Gly Thr Pro Phe Glu Thr Pro Asp Gin Gly Lys Ala
E--> 214
     215
E--> 216
           Arg Leu Leu Thr His Trp Glu/Gin Met Asp Tyr Gly Val (Gin )Phe Thr
     217
                            85
                                                 90
           Ala Ser Arg Lys Phe Leu Thr Ile Thr Pro Ile Val Leu Tyr Phe Leu
     218
     219
                       100
           Thr Ser Phe Tyr Thr Lys Tyr Asp(Gin) Ile His Phe Val Leu Asn Thr
E--> 220
                                        120
     221
E--> 222
           Val Ser Leu Met Ser Val Leu Ile Pro Lys Leu Pro Gin Leu His Gly
     223
                                    135
           Val Arg Ile Phe Gly Ile Asn Lys Tyr
     224
     225
     254 <210> SEQ ID NO: 7
     255 <211> LENGTH: 200
                                                                   use lower-case l,
le ASP Ala hot rumeral
     256 <212> TYPE: PRT
     257 <213> ORGANISM: Homo sapiens
     259 <400> SEQUENCE: 7
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E--> 260
                                                 10
     261
     262
           Lys Leu Arg Arg Pro Met Val Ile Glu Ile Glu Lys Asn Phe Asp
     263
                                             25
     264
           Tyr Leu Arg Lys Glu Met Thr Gln Asn Ile Tyr Gln Met Ala Thr Phe
     265
                                         40
     266
           Gly Thr Thr Ala Gly Phe Ser Gly Ile Phe Ser Asn Phe Leu Phe Arg
     267
     268
           Arg Cys Phe Lys Val Lys His Asp Ala Leu Lys Thr Tyr Ala Ser Leu
     269
                                70
     270
           Ala Thr Leu Pro Phe Leu Ser Thr Val Val Thr Asp Lys Leu Phe Val
     271
                            85
     272
           Ile Asp Ala Leu Tyr Ser Asp Asn Ile Ser Lys Glu Asn Cys Val Phe
     273
                                           105
           Arg Ser Ser Leu Ile Gly Ile Val Cys Gly Val Phe Tyr Pro Ser Ser
     274
     275
                                        120
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Leu Ala Phe Thr Lys Asn Gly Arg Leu Ala Thr Lys Tyr His Thr Val

276

Input Set: A:\(01997.025700) Sequence Listing.txt
Output Set: N:\CRF4\03192003\J019151.raw

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277
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                                                        140
     278
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                               150
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                                               170
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     638 <222> LOCATION: (71)...(1051)
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                                                                                60
     642
           gacgetgace atg gee aag atg gag etc teg aag gee tte tet gge eag
                                                                               109
     643
                      Met Ala Lys Met Glu Leu Ser Lys Ala Phe Ser Gly Gln
     644
     645
           cgg aca ctc cta tct gcc atc ctc agc atg cta tca ctc agc ttc tcc
                                                                               157
     646
           Arg Thr Leu Leu Ser Ala Ile Leu Ser Met Leu Ser Leu Ser Phe Ser
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                                    20
     648
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                                                                               205
     650
           Thr Thr Ser Leu Leu Ser Asn Tyr Trp Phe Val Gly Thr Gln Lys Val
     651
            30
     652
           ccc aag ccc ctg tgc gag aaa ggt ctg gca gcc aag tgc ttt gac atg
                                                                               253
     653
           Pro Lys Pro Leu Cys Glu Lys Gly Leu Ala Ala Lys Cys Phe Asp Met
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                            50
                                                55
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           cca gtg tcc ctg gat gga gat acc aca tcc acc cag gag gtg gta
                                                                               301
     656
           Pro Val Ser Leu Asp Gly Asp Thr Asn Thr Ser Thr Gln Glu Val Val
     657
                                            70
     658
           caa tac aac tgg gag act ggg gat gac cgg ttc tcc ttc cgg agc ttc
                                                                               349
     659
           Gln Tyr Asn Trp Glu Thr Gly Asp Asp Arg Phe Ser Phe Arg Ser Phe
     660
                                        85
     661
           cgg agt ggc atg tgg cta tcc tgt gag gaa act gtg gaa gaa cca ggg
                                                                               397
     662
           Arg Ser Gly Met Trp Leu Ser Cys Glu Glu Thr Val Glu Glu Pro Gly
     663
                                   100
     664
           gag agg tgc cga agt ttc att gaa ctt aca cca cca gcc aag aga gaa
     665
           Glu Arg Cys Arg Ser Phe Ile Glu Leu Thr Pro Pro Ala Lys Arg Glu
     666
           110
                               115
                                                    120
     667
           atc cta tgg tta tcc ctg gga acg cag atc acc tac atc gga ctt caa
W-->668
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     669
                          130
                                               135
     670
           ttc atc agc ttc ctc ctg cta cta aca gac ttg cta ctc act ggg aac
     671
           Phe Ile Ser Phe Leu Leu Leu Thr Asp Leu Leu Thr Gly Asn
     672
                                           150
     673
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RAW SEQUENCE LISTING

PATENT APPLICATION: US/10/019,151 TIME: 14:33:58

DATE: 03/19/2003

Input Set : A:\(01997.025700) Sequence Listing.txt
Output Set: N:\CRF4\03192003\J019151.raw

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676 ctg tca cs system for the component of the component		674	Pro	Ala	Cys	Gly	Leu	Lys	Leu	Ser	Ala	Phe	Ala	Ala	Val	Ser	Ser	Val		÷
N> 677 678 679 679 679 679 679 679 679 679 679 679		675			160					165					170					
N> 677 678 679 679 679 679 679 679 679 679 679 679		676	ctg	tca	P	ctc	ctg	ggg	atg	gtg	gcc	cac	atg	atg	tat	tca	caa	gtc	637	
678	W>	677																		
680 Phe Gln Ālā Thr Val Asn Leu Gly Pro Glu Āsp Trp Arg Pro His Val 681 190 205 682 tgg sat tat ggc tgg gcc ttc tac atg gcc tgc ctc ctc tcc acc tgc 683 Trp Asn Tyr Gly Trp Ala Phe Tyr Met Ala Trp Leu Ser Phe Thr Cys 684 210 215 220 685 tgc atg gcc tcg gcc tgc acc acc ttc aac acg tac acc agg atg gtg 686 Cys Met Ala Ser Ala Val Thr Thr Phe Asn Thr Tyr Thr Arg Met Val 687 225 230 688 ctg agg ttc aag tgc aag cat agt aag agc ttc aag gaa acc cg aac 689 Leu Glu Phe Lys Cys Lys His Ser Lys Ser Phe Lys Glu Asn Pro Asn 690 240 245 250 691 tgc cta cac act cac cat cag tgt ttc cct cgg cgg ctg ta agt ga ga 692 Cys Leu Pro His His His Gln Cys Phe Pro Arg Arg Leu Ser Ser Ala 693 255 260 265 694 gcc cca acc gtg ggt cct ttg acc agc tac cac cag tat cat aat cag 695 Ala Pro Thr Val Gly Pro Leu Thr Ser Tyr His Gln Tyr His Asn Gln 696 270 275 280 697 ccc atc cac tct gtc tct gac gag agg gct gct cta cag gag 698 Pro 11e His Ser Val Ser Glu Gly Val Asp Phe Tyr Ser Glu Leu Arg 699 290 290 300 690 8-700 aac aag gga de caa aga agg gcd gac cag aga aga aga gct 699 Arg Ser Ser Val Glu Glu Glu Gln Cys 700 305 310 315 701 Asn Lys Gly Phe Gln Arg Gly Ala Ser Gln Glu Leu Lys Glu Ala Val 702 305 320 325 706 agtagacttg agacctacct tacacgtctg ctgattatca acatgtgctt aagcc 708 <210 > SEO ID NO: 20 709 <211 > LENGTH: 1734 710 <212 > TyPE: DNA 711 <221 > NAME/KEY: CDS 715 <222 > LOCATION: (40) (1014) 717 <400 > SEQUENCE: 20 718 ctcttgagg gaccgtacct tacacgtctg ctgattatca acatgtgctt aagcc 710 cc gag aac aga cgc cgc gac at ggc gcc gc gcg gct 720 271 Cc gag aac aga cgc cgc gcc at cgg cc cgc gcc cgc gcc gcc gcc gcc gcc		678		175	$\sqrt{}$	<i>'</i>		_	180					185	_					
681 190 195 200 205 682 tgg aat tat ggc tgg gcc ttc tac at ag gcc tgg ctc tcc tcc acc tgc 683 Trp Asn Tyr Gly Trp Ala Phe Tyr Met Ala Trp Leu Ser Phe Thr Cys 684 210 215 220 685 tgc atg gct gcg gct gca acc acc tcc aac acc aga aga gtg gtg 686 Cys Met Ala Ser Ala Val Thr Thr Phe Asn Thr Tyr Thr Arg Met Val 687 225 230 235 688 ctg gag ttc aag tgc aag cat agt aag agc ttc aag gaa aac ccg aac 689 Leu Glu Phe Lys Cys Lys His Ser Lys Ser Phe Lys Glu Asn Pro Asn 690 240 245 250 691 tgc cta cca cat cac cat cag tgt ttc cct cgg cgg ctg tca agt gca 692 Cys Leu Pro His His His Gln Cys Phe Pro Arg Arg Leu Ser Ser Ala 693 255 260 265 694 gcc cca acc gtg ggt cct ttg acc agc tac cac cat tac acc at tgt ctga gg agg agg acc acc acc acc acc acc tcg ttc gag gga gtc gac ttc acc acc acc acc acc tcg tcc ttg ag gga gtc gac ttc acc acc acc acc acc act gtc ttc gag gga gtc gac ttc acc acc acc acc acc acc tcg tcc ttc gag gga gtc gac ttc acc acc aga gag 698 Pro Tle His Ser Val Ser Glu Gly Val Asp Phe Tyr Ser Glu Leu Arg 699 290 295 300 690 290 295 300 691 30 305 310 315 703 agg tca tct gta gag gaa gag cag tgt tagagattaa gcgggtttgg gg 704 Arg Ser Ser Val Glu Glu Gln Cys 705 320 325 706 agtaggettg agccctacct tacacgtctg ctgattatca acatgtgctt aagcc 708 <210 SEQ ID NO: 20 709 <211 SENGTH: 1734 710 <212 Type: DNA 711 <213 ORGANISM: Homo sapiens 713 <220 FEATURE: 714 <221 NAME/KSY: CDS 715 <222 LOCATION: (40)(1014) 717 <400 SEQUENCE: 20 718 ctcttggge gccggtggg ggcggcccg gcagggggg atg gcg gcc gcg gct 74		679	ttc	caa	gcg	act	gtc	aac	ttg	ggt	cca	gaa	gac	tgg	aga	cca	cat	gtt	685	
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RAW SEQUENCE LISTING

DATE: 03/19/2003 PATENT APPLICATION: US/10/019,151 TIME: 14:33:58

Input Set : A: $\(01997.025700)$ Sequence Listing.txt Output Set: N:\CRF4\03192003\J019151.raw

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	728	Asp	Leu	Ser	His	Arg	Asn	Lys	Glu	Pro	Pro	Ala	Pro	Ala	Gln	Gln	Leu		
	729			40					45					50					
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E>	733	aaa	ata	ttt	aca	cca	gca	gct	cca	gtt	cat	acc	aat	aaa	gaa	gat	qe)t		لمنامر 294
	734	Lys	Ile	Phe	Thr	Pro	Ala	Ala	Pro	Val	His	Thr	Asn	Lys	Glu	Asp	Pro		
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	740	Ser	Val	Ile		Val	Ser	Glu	Leu		Asp	Lys	Thr	Phe		Ile	Ala		
	741				105					110					115				
	742	_		_	_	_	_			_	_	acc		_	-		_		438
	743	Ala	Ile		Ala	Met	Arg	Tyr		Arg	Leu	Thr	Val		Ala	Gly	Ala		
	744			120					125					130					
	745											tca							486
	746	Met		Ala	Leu	GLy	Leu		Thr	Cys	Leu	Ser		Leu	Phe	GLy	Tyr		
	747		135					140					145						504
	748	_			-				-			tac		_			-		534
	749		Thr	Thr	vaı	тте		Arg	vaı	Tyr	Tnr	Tyr	Tyr	vaı	Ser	Tnr			
	750 751	150	+++	~~~	-++	+++	155	-++	. ~ .	a+~	a++	160	~~~	~~~	++-		165		E02
	752											cgg Arg							582
	753	ьец	rne	Ата	116	170	сту	116	ALG	Met	175	Arg	GIU	СТУ	neu	180	Mec		
	754	age	cct	rat	nan		caa	nan	ma a	cta	-	gaa	att	C22	act		tta		630
	755											Glu							050
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	764	Leu	His	Phe	·Ile	Ser	Pro	Ile	Phe	Val	Gln	Ala	Leu	Thr	Leu	Thr	Phe		
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	766											act							822
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Input Set : A:\(01997.025700) Sequence Listing.txt
Output Set: N:\CRF4\03192003\J019151.raw

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777		295					300					305					
778					tct												1011
779	Ala	Phe	Ala	Phe	Ser	Ala	Leu	Phe	Ile	Ser	Pro	Asp	Ser	Gly	Phe		
780	310					315					320						
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790	ctg	ccag	gtc a	attc	ttcct	c ti	tttt	tttt	aat	tgg	gtag	gaca	accca	aat a	ataaa	aaca	g 1610
791	tcaa	atatt	tg a	acaat	tgtg	ga at	taco	caaat	taa	aaaga	agaa	tact	tatga	aat o	gtatt	tcatat	1670
792	ttt	tcta	ata t	tgaa	ataaa	ac aa	atgta	aacat	aga	ataad	caat	ataa	aataa	aaa (gtggt	tatgad	1730
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979	<222>	TOO	·	T. / [- O \	(62	771					11	/ 1				
						(02	41)					V	1				
981	<400>	SEQ	JENCE	E: 24	4		·					V					
981 982	<400> aca	SEQU caca	JENCE cat o	E: 24 ctgca	4 accto	ca a	ccaca	-				aact	tggc1		tggg		57
981 982 983	<400> acad atg	SEQU cacad agg	JENCE cat of ctg	E: 24 ctgca tca	4 accto ctg	ca ao	ccaca ctg	ctg	ctg	ctg	ctg	aact ctg	tggci gga	gcc	tgg	gcc	57 105
981 982 983 984	<400> acad atg Met	SEQU cacad agg	JENCE cat of ctg	E: 24 ctgca tca	accto ctg Leu	ca ao	ccaca ctg	ctg	ctg	ctg Leu	ctg	aact ctg	tggci gga	gcc	tgg Trp	gcc	
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981 982 983 984 985 986 987 988 990 991	<400> acad atg Met 1 atc Ile caa Gln cgc	SEQUE ACA ATG CCA Pro Ctg Leu	JENCE cat of ctg Leu ggg Gly gat Asp 35 gat	E: 24 tca tca Ser ggc Gly 20 gat Asp	decctor ctg Leu 5 ctc Leu gag Glu tgc	ca acca cca Pro ggg Gly gag Glu aga	ccaca ctg Leu gac Asp atg Met	ctg Leu agg Arg tac Tyr 40 gtg	ctg Leu gcg Ala 25 tca Ser	ctg Leu 10 cca Pro gcc Ala	ctg Leu ctc Leu cac His	aact ctg Leu aca Thr atg Met	tggct gga Gly gcc Ala ccc Pro 45	gcc Ala aca Thr 30 gct Ala	tgg Trp 15 gcc Ala cac His	gcc Ala cca Pro ctg Leu	105 153
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981 982 983 984 985 986 987 988 990 991 992 993	<400> acad atg Met 1 atc Ile caa Gln cgc Arg	SEQUENCE AGE AGE AGE AT A CCA Pro Ctg Leu tgt Cys 50	JENCE cat of ctg Leu ggg Gly gat Asp gat Asp	etgea tea Ser ggc Gly 20 gat Asp gcc Ala	dector ctg Leu 5 ctc Leu gag Glu tgc Cys	ca ac cca Pro ggg Gly gag Glu aga Arg	ccaca ctg Leu gac Asp atg Met gct Ala 55	ctg Leu agg Arg tac Tyr 40 gtg Val	ctg Leu gcg Ala 25 tca Ser gct Ala	ctg Leu 10 cca Pro gcc Ala tac Tyr	ctg Leu ctc Leu cac His cag Gln	aact ctg Leu aca Thr atg Met atg Met 60	tggct gga Gly gcc Ala ccc Pro 45 tgg	gcc Ala aca Thr 30 gct Ala caa Gln	tgg Trp 15 gcc Ala cac His aat Asn	gcc Ala cca Pro ctg Leu ctg Leu	105 153 201 249
981 982 983 984 985 986 987 988 999 991 992 993 994 995	<400> acad atg Met 1 atc Ile caa Gln cgc Arg	SEQUENCE AGE AGE AGE AGE AGE AGE AGE AGE AGE AG	JENCE cat of ctg Leu ggg Gly gat Asp 35 gat Asp	etgea tea Ser ggc Gly 20 gat Asp gcc Ala	dector ctg Leu 5 ctc Leu gag Glu tgc Cys	ca ac cca Pro ggg Gly gag Glu aga Arg	ccaca ctg Leu gac Asp atg Met gct Ala 55 ctt	ctg Leu agg Arg tac Tyr 40 gtg Val	ctg Leu gcg Ala 25 tca Ser gct Ala acc	ctg Leu 10 cca Pro gcc Ala tac Tyr	ctg Leu ctc Leu cac His cag Gln	aact ctg Leu aca Thr atg Met atg Met 60 tct	tggct gga Gly gcc Ala ccc Pro 45 tgg Trp	gcc Ala aca Thr 30 gct Ala caa Gln	tgg Trp 15 gcc Ala cac His aat Asn	gcc Ala cca Pro ctg Leu ctg Leu cgg	105 153 201
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981 982 983 984 985 986 987 988 990 991 992 993 994 995 996 997 998	<400> acac atg Met 1 atc Ile caa Gln cgc Arg gca Ala 65 gag Glu	sequence agg Arg cca Pro ctg Leu tgt Cys aag Lys ctg Leu	JENCE cat of ctg Leu ggg Gly gat Asp gat Asp gca Ala agc Ser	tca ser ggc Gly 20 gat Asp gcc Ala gag Glu gag	dector ctg Leu 5 ctc Leu gag Glu tgc Cys acc Thr ttg Leu 85	ggg Gly gag Glu aga Arg aaa Lys 70 gtc Val	gac Asp atg Met State Leu tac Tyr	ctg Leu agg Arg tac Tyr 40 gtg Val cat His acg Thr	ctg Leu gcg Ala 25 tca Ser gct Ala acc Thr	Leu 10 cca Pro gcc Ala tac Tyr tca Ser gtc Val 90	ctg Leu ctc Leu cac His cag Gln aac Asn 75 ctg Leu	aact ctg Leu aca Thr atg Met 60 tct Ser gac	gga Gly gcc Ala ccc Pro 45 tgg Trp ggg Gly	gcc Ala aca Thr 30 gct Ala caa Gln ggg Gly agc ser	tgg Trp 15 gcc Ala cac His aat Asn cgg Arg tgc	gcc Ala cca Pro ctg Leu ctg Leu cgg Arg 80 tcc Ser 95	105 153 201 249 297 345
981 982 983 984 985 986 987 988 990 991 992 993 994 995 997 998 999	<400> acac atg Met 1 atc Ile caa Gln cgc Arg gca Ala 65 gag Glu 0 cgg	SEQUENCE AS A Process of the Control	JENCE cat of cat	tca ser ggc Gly 20 gat Asp gcc Ala gag Glu gag	dacctor ctg Leu 5 ctc Leu gag Glu tgc Cys acc Thr ttg Leu 85 gag	ggg Gly gag Glu aga Arg aaa Lys 70 gtc Val	gac Asp atg Met S55 ctt Leu tac Tyr	Leu agg Arg tac Tyr 40 gtg Val cat His acg Thr	ctg Leu gcg Ala 25 tca Ser gct Ala acc Thr gat Asp	Leu 10 cca Pro gcc Ala tac Tyr tca Ser gtc Val 90 a gaa	ctg Leu ctc Leu cac His cag Gln aac Asn 75 ctg Leu	aact ctg Leu aca Thr atg Met 60 tct Ser gac	gga Gly gcc Ala ccc Pro 45 tgg Trp ggg Gly	aca Thr 30 gct Ala caa Gln ggg Gly agc ser	tgg Trp 15 gcc Ala cac His aat Asn cgg Arg tgc Cys	cca Pro ctg Leu ctg Leu cgg Arg 80 tcc Ser	105 153 201 249
981 982 983 984 985 986 987 988 990 991 992 993 994 995 996 997 998	<400> acac atg Met 1 atc Ile caa Gln cgc Arg gca Ala 65 gag Glu 0 1 cgg 2 Arg	SEQUENCE AND AND ADDRESS OF THE ADDR	JENCE cat of cat	tca ser ggc Gly 20 gat Asp gcc Ala gag Glu gag	dacctor ctg Leu 5 ctc Leu gag Glu tgc Cys acc Thr ttg Leu gag	ggg Gly gag Glu aga Arg aaa Lys 70 gtc Val	gac Asp atg Met S55 ctt Leu tac Tyr	Leu agg Arg tac Tyr 40 gtg Val cat His acg Thr	ctg Leu gcg Ala 25 tca Ser gct Ala acc Thr gat Asp	Leu 10 cca Pro gcc Ala tac Tyr tca Ser gtc Val gas Gas Gli	ctg Leu ctc Leu cac His cag Gln aac Asn 75 ctg Leu	aact ctg Leu aca Thr atg Met 60 tct Ser gac	gga Gly gcc Ala ccc Pro 45 tgg Trp ggg Gly	aca Thr 30 gct Ala caa Gln ggg Gly agc ser	tgg Trp 15 gcc Ala cac His aat Asn cgg Arg tgc Cys	gcc Ala cca Pro ctg Leu ctg Leu cgg Arg 80 tcc Ser 95	105 153 201 249 297 345

Input Set : A:\(01997.025700) Sequence Listing.txt
Output Set: N:\CRF4\03192003\J019151.raw

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	1009		130		_	_		135			_		140	-		_		
	1010	cac	tac	ttg	ggg	gag	ttt	gga	gaa	gac	cag	atc	tat	gaa	gcc	cac	caa	537
	1011	His	Tyr	Leu	Gly	Glu	Phe	Gly	Glu	Asp	Gln	Ile	Tyr	Glu	Ala	His	Gln	
	1012	145	_		_		150·	_		_		155	_				160	
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	1014															Gln		
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	1017	_					Val		-		-	-			_			
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	1020	gcad	ggca	qcc a	aggad	gggc	ag ga	aaqc	cctt	g cto	ctat	acta	cca.	tcct	gcc ·	tccci	tcctcc	750
	1021	-		-				-		_					_		aataaa	810
	1022	_		gac (- , , , , ,		- 2 2 2		,					ر ر			823
E>				322			A 0	1										020
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<400> 17 cttgcctctg ggaaggaaat acattataga gtgggaaatt tttatcattt tgaaccaaga 60 ttcttctaaa gaaagaaaga ctgattaata aaatgtggca gctgtgctct tcaaggcatt 120 180 tatagtgtat atagttttag aaaaacagtc ccaccactta agcatagatg taatttacta ataaaa atg att ctg ctt gtg att ctt gca ttt tat ctg tgg cag gtg 228 Met Ile Leu Leu Val Ile Leu Ala Phe Tyr Leu Trp Gln Val 5 10 gac atg ttg agt gaa att aac att gct ccc cgg att ctc acc aat ttc 276 Asp Met Leu Ser Glu Ile Asn Ile Ala Pro Arg Ile Leu Thr Asn Phe 25 20 act gga gta atg cca cct cag ttc aaa aag gat ttg gat tcc tat ctt 324 Thr Gly Val Met Pro Pro Gln Phe Lys Lys Asp Leu Asp Ser Tyr Leu 35 35-40 aaa act cga tca cca gtc act ttc ctg tct gat ctg cgc agc aac cta 372 Lys Thr Arg Ser Pro Val Thr Phe Leu Ser Asp Leu Arg Ser Asn Leu Please covert all misabered among number throughout Sequence Listing

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VERIFICATION SUMMARY

PATENT APPLICATION: US/10/019,151

Input Set : A:\(01997.025700) Sequence Listing.txt
Output Set: N:\CRF4\03192003\J019151.raw

DATE: 03/19/2003

TIME: 14:33:59

L:11 M:270 C: Current Application Number differs, Replaced Current Application No L:11 M:271 C: Current Filing Date differs, Replaced Current Filing Date L:27 M:332 E: (32) Invalid/Missing Amino Acid Numbering, SEQ ID:1 M:332 Repeated in SeqNo=1 L:82 M:332 E: (32) Invalid/Missing Amino Acid Numbering, SEQ ID:2 M:332 Repeated in SeqNo=2 L:147 M:330 E: (2) Invalid Amino Acid Designator, NUMBER OF INVALID KEYS:1 L:214 M:330 E: (2) Invalid Amino Acid Designator, NUMBER OF INVALID KEYS:1 L:216 M:330 E: (2) Invalid Amino Acid Designator, NUMBER OF INVALID KEYS:2 L:220 M:330 E: (2) Invalid Amino Acid Designator, NUMBER OF INVALID KEYS:1 L:222 M:330 E: (2) Invalid Amino Acid Designator, NUMBER OF INVALID KEYS:1. L:260 M:330 E: (2) Invalid Amino Acid Designator, NUMBER OF INVALID KEYS:1 L:494 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:497 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:500 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:503 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:506 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:509 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:512 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:515 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:518 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:521 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:524 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:527 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:530 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:17 L:574 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:576 M:334 W: (2) Invalid Amino Acid in Coding Region, NUMBER OF INVALID KEYS:1 L:577 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:579 M:334 W: (2) Invalid Amino Acid in Coding Region, NUMBER OF INVALID KEYS:1 L:580 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:583 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:586 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:588 M:334 W: (2) Invalid Amino Acid in Coding Region, NUMBER OF INVALID KEYS:1 L:589 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:592 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:594 M:334 W: (2) Invalid Amino Acid in Coding Region, NUMBER OF INVALID KEYS:1 L:595 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:598 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:601 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:604 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:607 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:610 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:613 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:617 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:18 L:668 M:334 W: (2) Invalid Amino Acid in Coding Region, NUMBER OF INVALID KEYS:1 L:677 M:334 W: (2) Invalid Amino Acid in Coding Region, NUMBER OF INVALID KEYS:1 L:700 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:1

L:733 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:1

VERIFICATION SUMMARY

PATENT APPLICATION: US/10/019,151

DATE: 03/19/2003 TIME: 14:33:59

Input Set : A:\(01997.025700) Sequence Listing.txt

Output Set: N:\CRF4\03192003\J019151.raw

L:825 M:334 W: (2) Invalid Amino Acid in Coding Region, NUMBER OF INVALID KEYS:1 L:999 M:334 W: (2) Invalid Amino Acid in Coding Region, NUMBER OF INVALID KEYS:1

L:1015 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:24 L:1018 M:336 W: Invalid Amino Acid Number in Coding Region, SEQ ID:24

L:1024 M:258 W: Mandatory Feature missing, <223> Tag not found for SEQ ID#:24

L:1024 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:24 after pos.:823 L:1024 M:254 E: No. of Bases conflict, LENGTH:Input:1 Counted:831 SEQ:24

L:1024 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:8

L:1024 M:112 C: (48) String data converted to lower case,

L:1024 M:252 E: No. of Seq. differs, <211> LENGTH:Input:823 Found:831 SEQ:24